PCT/JP2007/063946

#### From the INTERNATIONAL BUREAU

NOTIFICATION OF TRANSMITTAL OF COPIES OF TRANSLATION OF THE INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (CHAPTER I OR CHAPTER II OF THE PATENT COOPERATION TREATY)

(PCT Rules 44bis.3(c) and 72.2)

To:

SHIMIZU, Hatsushi Kantetsu Tsukuba Bldg. 6F 1-1-1, Oroshi-machi Tsuchiura-shi, Ibaraki 3000847 **JAPON** 



Date of mailing (day/month/year) 29 January 2009 (29.01.2009)

Applicant's or agent's file reference C1-A0604P

International application No. PCT/JP2007/063946 IMPORTANT NOTIFICATION

International filing date (day/month/year) 13 July 2007 (13.07.2007)

Applicant

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CHUGAI SEIYAKU KABUSHIKI KAISHA et al

١.	Transmittal of	the	translation	to	the	applicant.
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The International Bureau transmit patentability (Chapter I).	herewith a copy of the English	th translation of the internation	nal preliminary report or
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The International Bureau transmits herewith a copy of the English translation of the international preliminary report on patentability (Chapter II).

#### Transmittal of the copy of the translation to the designated or elected Offices.

The International Bureau notifies the applicant that copies of that translation have been transmitted to the following designated or elected Offices requiring such translation:

The following designated or elected Offices, having waived the requirement for such a transmittal at this time, will receive copies of that translation from the International Bureau only upon their request:

AE, AG, AL, AM, AP, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EA, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OA, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM,

Reminder regarding translation into (one of) the official language(s) of the elected Office(s).

The applicant is reminded that, where a translation of the international application must be furnished to an elected Office, that translation roust contain a translation of any annexes to the international preliminary report on patentability (Chapter II).

It is the applicant's responsibility to prepare and furnish such translation directly to each elected Office concerned within the applicable time limit (Rule 74.1). See Volume II of the PCT Applicant's Guide for further details.

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

Authorized officer

Yoshiko Kuwahara

e-mail: pt07.pct@wipo.int

Facsimile No. +41 22 338 82 70

### PATENT COOPERATION TREATY

# **PCT**

### INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter I of the Patent Cooperation Treaty)

(PCT Rule 44bis)

Applicant's or agent's file reference C1-A0604P	FOR FURTHER ACTION	See item 4 below			
International application No. PCT/JP2007/063946	International filing date (day/month/year) 13 July 2007 (13.07.2007)	Priority date (day/month/year) 13 July 2006 (13.07.2006)			
International Patent Classification (8th edition unless older edition indicated) See relevant information in Form PCT/ISA/237					
Applicant CHUGAI SEIYAKU KABUSHIKI KAISHA					

1.	This international preliminary International Searching Autho	report on patentability (Chapter I) is issued by the International Bureau on behalf of the rity under Rule 44 bis.1(a).
2.	This REPORT consists of a to	tal of 9 sheets, including this cover sheet.
		rence to the written opinion of the International Searching Authority should be read as a reference y report on patentability (Chapter I) instead.
3.	This report contains indication	s relating to the following items:
	Box No. I	Basis of the report
	Box No. II	Priority
	Box No. III	Non-establishment of opinion with regard to novelty, inventive step and industrial applicability
	Box No. IV	Lack of unity of invention
	Box No. V	Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement
	Box No. VI	Certain documents cited
	Box No. VII	Certain defects in the international application
	Box No. VIII	Certain observations on the international application
4.		communicate this report to designated Offices in accordance with Rules 44bis.3(c) and 93bis.1 but makes an express request under Article 23(2), before the expiration of 30 months from the priority

	Date of issuance of this report 20 January 2009 (20.01.2009)
The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland	Authorized officer Yoshiko Kuwahara
Facsimile No. +41 22 338 82 70	e-mail: pt07.pct@wipo.int

PATENT COOPERATION TREATY

From th		NAL SEARCHIN	IG AUTHORI	ITY			RANG.
To:							PCT PCT
					INTE		EN OPINION OF THE AL SEARCHING AUTHORITY
						(P	PCT Rule 43bis.1)
					Date of mail	_	
ł		gent's file referen	re		FOR FURT	THER ACTIO	
	-A060		<del></del> ,		<u> </u>		aragraph 2 below
	-	plication No. 2007/063:	I	International filing date 13.07.2007	(day/month/yea		ity date (day/month/year) 3.07.2006
Internat	tional Pa	tent Classification	(IPC) or both	national classification ar	nd IPC		
Applica	nt						
		SEIYAKU	KABUSH	IKI KAISHA			
1.	This o	Dinion contains in	dications relati	ing to the following item	s:		
	$\boxtimes$	Box No. I	Basis of the o				
	Box No. II Priority						
	Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability						
	Box No. IV Lack of unity of invention						
	Box No. V  Reasoned statement under Rule 43bis.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement						
	Box No. VI Certain documents cited						
	Box No. VII Certain defects in the international application						
	$\bowtie$	Box No. VIII	Certain obser	evations on the internation	nal application		
2.	FURT	HER ACTION					
	Internation than the	ational Preliminar his one to be the I	y Examining A PEA and the c	authority ("IPEA") excep	t that this does the Internation	not apply whe	considered to be a written opinion of the effect of the applicant chooses an Authority of the Rule $66.1bis(b)$ that written opinions
	writter	reply together.	where appropr		before the exp	iration of 3 i	applicant is invited to submit to the IPEA months from the date of mailing of Fo states.
	For fur	ther options, see	Form PCT/ISA	/220.			
3.	For fur	ther details, see n	otes to Form P	CT/ISA/220.			
Name a	nd maili	ng address of the I	SA/JP	Date of completion of	of this opinion	Authorized	officer
		-			•		
Facsimi	le No.					Telephone N	۷٥.

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	EVICENTIONAL SEARCHENG ACTION II	FC1/0F2007/003940
Box N	o. I Basis of this opinion	
1. 1	With regard to the language, this opinion has been established on the basis of:	
	the international application in the language in which it was filed	
	the translation of the international application into	, which is the language of a
	translation furnished for the purposes of international search (Rule 12.3(a) and 23.1(	b)).
i	With regard to any nucleotide and/or amino acid sequence disclosed in the internancention, this opinion has been established on the basis of:  a. type of material	tional application and necessary to the claimed
	a sequence listing	
	table(s) related to the sequence listing	
	b. format of material	
	on paper	
	in electronic form	
	c. time of filing/furnishing	
	contained in the international application as filed	
	filed together with the international application in electronic form	
	furnished subsequently to this Authority for the purposes of search	
3. L	In addition, in the case that more than one version or copy of a sequence listing furnished, the required statements that the information in the subsequent or addition filed or does not go beyond the application as filed, as appropriate, were furnished.	
<b>4</b> . A	additional comments:	
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### WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No.

PCT/JP2007/063946 Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability The questions whether the claimed invention appears to be novel, to involve an inventive step (to be non obvious), or to be industrially applicable have not been examined in respect of: the entire international application claims Nos. the said international application, or the said claims Nos. subject matter which does not require an international search (specify): the description, claims or drawings (indicate particular elements below) or said claims Nos. 8 are so unclear that no meaningful opinion could be formed (specify): Claim 8 specifies an antibody with a function of "binding to an epitope the same as that of a human leukocyte antigen (HLA) protein to which the antibody in any one of claims 1-7 binds", but no epitope recognized by the antibody in the invention of this application is specified in the description of this application, and therefore claim 8 is quite obscure because it is not clear what antibody is encompassed. the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed (specify): no international search report has been established for said claims Nos. 8 a meaningful opinion could not be formed without the sequence listing; the applicant did not, within the prescribed time limit: furnish a sequence listing on paper complying with the standard provided for in Annex C of the Administrative Instructions, and such listing was not available to the International Searching Authority in a form and manner acceptable to it. furnish a sequence listing in electronic form complying with the standard provided for in Annex C of the Administrative Instructions, and such listing was not available to the International Searching Authority in a form and manner acceptable to it. pay the required late furnishing fee for the furnishing of a sequence listing in response to an invitation under Rules 13ter.1(a) a meaningful opinion could not be formed without the tables related to the sequence listings; the applicant did not, within the prescribed time limit, furnish such tables in electronic form complying with the technical requirements provided for in Annex C-bis of the Administrative Instructions, and such tables were not available to the International Searching Authority in a form and manner the tables related to the nucleotide and/or amino acid sequence listing, if in electronic form only, do not comply with the technical requirements provided for in Annex C-bis of the Administrative Instructions. See Supplemental Box for further details.

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	citations and expla	nations supporting such statement	
1.	Statement		
	Novelty (N)	Claims 1-7, 9-25	YES
		Claims	NO
	Inventive step (IS)	Claims 1-7, 9-25	YES
		Claims	NO NO
	Industrial applicability (IA)	Claims 1-7, 9-25	YES
		Claims	NO

Reasoned statement under Rule 43bis. I(a)(i) with regard to novelty, inventive step or industrial applicability;

#### 2. Citations and explanations:

Box No. V

Stanned

- Document 1: WO 2004/033499 A1 (Chugai Pharmaceutical Co., Ltd.), 22 April 2004, full text & EP 1561759 A1
- Document 2: Kimura N. et al., 2D7 diabody bound to the alpha2 domain of HLA class I efficiently induces caspase-independent cell death against malignant and activated lymphoid cells, Biochem. Biophys. Res. Commun., 2004, Vol. 325, No. 4, pages 1201 to 1209, abstract
- Document 3: Genestier L. et al., Fas-independent apoptosis of activated T cells induced by antibodies to the HLA class I alphal domain, Blood, 1997, Vol. 90, No. 9, pages 3629 to 3639, abstract

Concerning claims 1-7 and 9-23

The invention as set forth in claims 1-7 and 9-23 is novel and involves an inventive step in view of documents 1-3 cited in the ISR.

Documents 1 and 2 describe that a monoclonal antibody to the  $\alpha 2$  domain of HLA was obtained, that a low-molecular antibody was fabricated using a variable region of said antibody and that the low-molecular antibody is cytotoxic.

Document 3 describes that an antibody to the lpha 2 domain of HLA was obtained and that said antibody inhibits cell

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Box No. V

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Reasoned statement under Rule 43bis.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

proliferation but does not cause apoptosis.

The antibody of the invention of this application is an antibody which specifically recognizes the  $\alpha 2$  domain of HLA, but there is no technique wherein CDRs 1, 2 and 3 of a heavy chain variable region of the anti-HLA  $\alpha 2$  domain antibody are identified with SEQ ID NOs: 7, 8 and 9, respectively, and CDRs 1, 2 and 3 of a light chain variable region are identified with SEQ ID NOs: 10, 11 and 12, respectively, and said antibody is a novel protein. Furthermore, it is apparent that the low-molecular antibody of the invention of this application has an excellent cell proliferation inhibiting capability as compared to the antibodies described in documents 1-3.

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Box No. VIII

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Certain observations on the international application

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made:

In the antibody in claims 1 and 3, the positions of CDR in the heavy chain variable region to which SEQ ID NOs: 7, 8 and 9 correspond are not adequately identified, and CDR1=SEQ ID NO: 7, CDR2=SEQ ID NO: 8 and CDR3=SEQ ID NO: 9 are merely mentioned in the description of the present application, and so the feature of the invention as in claims 1 and 3 is not considered to be set forth adequately in the description.

In the antibody in claims 2 and 3, the positions of CDR in the light chain variable region to which SEQ ID NOs: 10, 11 and 12 correspond are not adequately identified, and CDR1=SEQ ID NO: 10, CDR2=SEQ ID NO: 11 and CDR3=SEQ ID NO: 12 are merely mentioned in the description of the present application, and so the feature of the invention as in claims 2 and 3 is not considered to be set forth adequately in the description.

Concerning the antibody having "amino acid sequences with one or more amino acid sequences replaced, lost, inserted and/or added" in claims 4-7 (b) and (f), SEQ ID NOs: 2, 4 and 6 each include a CDR region, and if amino acid residues are "replaced, lost, inserted and/or added" in said region, the activity of said antibody is likely reduced, and whether a desired effect is exhibited is unclear.

Claim 25 claims "an autoimmune disease therapeutic agent containing as an effective ingredient the antibody set forth in any of claims 1-14", but it is not described that the disease could be actually treated using the antibody of the invention of this application, and so the feature of the

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			101/012001/003310	
Box No. VIII	Certain observations on the international appl	lication		
invent	ion as in claim 25 is a	not considered	to be set forth	
ì	tely in the description		to be set forth	
				i

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#### Supplemental Box

In case the space in any of the preceding boxes is not sufficient.

C12P21/08(2006.01)i

Continuation of: International Patent Classification (IPC) C12N15/00(2006.01)i, A61K39/395(2006.01)i, A61P7/00(2006.01)i, A61P35/00(2006.01)i, C07K16/18(2006.01)i, C12N1/15(2006.01)i, C12N1/19(2006.01)i, C12N1/21(2006.01)i, C12N5/10(2006.01)i, C12N15/09(2006.01)i,